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REGULAR ARTICLE

Genetic characterization of the Pss region and the role of PssS in exopolysaccharide production and symbiosis of *Rhizobium leguminosarum* bv. *trifolii* with clover

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Abstract

Background and aims In the symbiotic bacterium *Rhizobium leguminosarum* bv. *trifolii*, a majority of proteins involved in exopolysaccharide (EPS) synthesis are encoded by genes located in a large polysaccharide synthesis cluster (Pss). The aim of this study was genetic characterization of the Pss region in the Rt24.2 strain in the context of EPS production and symbiosis with red clover (*Trifolium pratense*).

Methods The expression of genes located in the Pss cluster was determined using constructed *pss-lacZ* transcriptional fusions. The role of transcriptional regulator RosR in *pss* transcription was confirmed using a *rosR* mutant and the Rt24.2(pBR1) strain carrying multiple *rosR* copies. An EPS-deficient mutant, Rt770 was obtained using a random mutagenesis and mTn5SSgusA40 transposon. Symbiotic properties of the Rt770 strain in interaction with clover were characterized in inoculation experiments. Infection of host roots and nodule

occupancy by this mutant were investigated using both light and electron microscopy.

Results Transcriptional levels of particular *pss* genes differed significantly; the genes encoding glycosyltransferases and enzymes modifying EPS have promoters of weak activities, whereas those encoding proteins involved in EPS polymerization and export possess stronger promoters. Furthermore, RosR affected expression of some *pss* genes. A mutation in Rt24.2 *pssS* encoding glucosyltransferase totally abolished EPS synthesis, decreased motility, and increased sensitivity to some stressors. The *pssS* mutant Rt770 induced formation of nodules on clover roots, which were ineffective in nitrogen fixation.

Conclusion EPS secreted by Rt24.2 is required for both adaptation to soil conditions and the establishment of effective symbiosis with clover plants.

Keywords *pss* genes · Transcription · *Rhizobium leguminosarum* · Exopolysaccharide synthesis · Glycosyltransferase PssS · Legume-rhizobium symbiosis

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Introduction

Extracellular polysaccharides (EPS) are produced by a wide range of microorganisms. Various functions are ascribed to these polymers, among them protection against stress factors and plant antimicrobial compounds, nutrient gathering, attachment to both abiotic and biotic surfaces, and biofilm formation, which

ensures adaptation of bacterial cells to changing environmental conditions (Fujishige et al. 2006; Flemming and Wingender 2010; Jaszek et al. 2014; Janczarek et al. 2015a). In the case of soil nitrogen-fixing bacteria, which are able to establish symbiotic interactions with legumes, EPS is also indispensable for effective infection and nodulation of many host plants (e.g., *Medicago*, *Trifolium*, *Pisum*, *Vicia* spp.) (Downie 2010; Janczarek et al. 2015b). Strains defective in EPS production from several rhizobial species, such as *S. meliloti*, and *Rhizobium leguminosarum* bvs. *trifolii* and *viciae*, induce only small, partially infected or even empty nodule-like structures on their host plants that are ineffective in nitrogen fixation (Cheng and Walker 1998; van Workum et al. 1998; Rolfe et al. 1996; Janczarek and Rachwał 2013).

R. leguminosarum bv. *trifolii* induces formation of nodules on clover roots (*Trifolium pratense*). The structure of EPS synthesized by this bacterium has been established in detail. It is a heteropolymer composed of octasaccharide repeating units consisting of a backbone of two glucose and two glucuronic acid residues, and a side chain of three glucose and one galactose residues. These subunits are additionally modified by non-sugar substituents (one *O*-acetyl and two pyruvyl groups) (Robertsen et al. 1981; McNeil et al. 1986; Cremers et al. 1991; O'Neill et al. 1991; Breedveld et al. 1993). However, the biosynthesis pathway of this polysaccharide in *R. leguminosarum* is known only fragmentarily. So far, functions of enzymes engaged in the four steps of the EPS unit assembly have been experimentally confirmed. The initiation of EPS biosynthesis is conducted by a glucosyl-IP-transferase, which transfers glucose-1-phosphate from UDP-glucose to a C₅₅-isoprenylphosphate (IP) carrier anchored at the cytoplasmic site of the inner membrane (Pollock et al. 1998). This enzyme is encoded by the *pssA* gene, which is located in the chromosome at a long distance from other EPS synthesis genes and transcribed as a monocistronic mRNA (Borthakur et al. 1988; Ivashina et al. 1994; van Workum et al. 1997). Glucuronosyl-(β 1-4)-glucosyl transferase PssDE and glucuronosyl-(β 1-4)-glucuronosyl transferase PssC are involved in the second and the third step of the unit synthesis, respectively (Pollock et al. 1998). These proteins are encoded by the *pssC*, *pssD*, and *pssE* genes located in a cluster named Pss-I (Król et al. 2007). Mutations in the *pssA* and *pssD* genes totally abolished EPS synthesis (Borthakur et al. 1988; Ivashina et al.

1994; Janczarek and Urbanik-Sypniewska 2013), whereas a disruption of the *pssC* gene resulted in a decrease in EPS synthesis to 40 % of the amount produced by the wild-type strain (van Workum et al. 1997). *pssJ* encodes a galactosyltransferase, which probably participates in the last step of the unit assembly (Breedveld et al. 1993). A strain *exo344::Tn5* carrying a mutation in the *pssJ* gene produced only residual amounts of EPS, whose repeating units did not contain terminal galactose.

However, genes involved in the remaining steps of the unit synthesis have not yet been identified, except *pssM*, which is located in this cluster. PssM is a ketal pyruvate transferase responsible for one of the non-sugar modifications of EPS (pyruvylation of the subterminal glucose in the repeating unit) (Ivashina et al. 2010). A mutation in *pssM* resulted in failed symbiosis with the compatible host plant. Moreover, some genes involved in EPS polymerization and export were characterized (*pssTNOP*, *pssL* and *pssP2*), among them a majority are located in the polysaccharide synthesis region (Mazur et al. 2005; Marczak et al. 2013, 2014).

The amount of produced EPS and its biochemical properties seem to be very important for functions of this polysaccharide in both adaptation to soil conditions and symbiosis. However, the knowledge of EPS biosynthesis in *R. leguminosarum* and factors engaged in the regulation of this process is still limited. Up to now, functions of four proteins (P_{si}A, P_{sr}A, ExoR, and RosR) have been confirmed. *psrA* and *psiA*, which are located on symbiotic megaplasmids of *R. leguminosarum* bv. *phaseoli* strains, were discovered as the first regulatory genes (Borthakur and Johnston 1987; Borthakur et al. 1988). Although a mutation in *psiA* did not affect the amounts of produced EPS, multiple copies of this gene prevented EPS synthesis. This effect was overcome in the presence of additional copies of *psrA* or *pssA*, indicating that balanced copy numbers of these genes are needed for a proper level of EPS production (Borthakur and Johnston 1987; Latchford et al. 1991; Mimmack et al. 1994). Another gene, *exoR* also negatively influenced EPS synthesis in *R. leguminosarum*, since an *exoR* mutant produced significantly more EPS than the wild-type strain (Reeve et al. 1997).

In contrast, a protein encoded by *rosR* positively affected EPS synthesis in *R. leguminosarum*. A mutation in this gene resulted in a 3-fold decrease, whereas multiple *rosR* copies increased the synthesis of this

polymer nearly 2-fold (Janczarek and Skorupska 2007; Janczarek et al. 2009). RosR belongs to the family of Ros/MucR transcriptional regulators, which are involved in regulation of EPS synthesis in several rhizobial species (*S. meliloti*, *Rhizobium etli*, *Agrobacterium tumefaciens*, *Agrobacterium radiobacter*) (D'Souza-Ault et al. 1993; Keller et al. 1995; Bittinger et al. 1997; Chou et al. 1998). This protein contains a Cys₂His₂-type zinc-finger motif responsible for binding to a 22-bp-long sequence called the RosR-box. Previously, it was confirmed that RosR affected transcription of *pssA* and its own gene (Janczarek and Skorupska 2007; Janczarek and Urbanik-Sypniewska 2013).

Up to now, no comprehensive analysis of transcription of genes involved in *R. leguminosarum* EPS synthesis has been performed so far. In this work, genetic organization of the polysaccharide synthesis cluster (Pss) of the Rt24.2 strain was established and a detailed transcriptional analysis of the *pss* genes located in it was performed. In addition, a mutant strain in one of the *pss* genes of this cluster (*pssS*) was obtained and characterized in relation to EPS production and symbiosis with clover.

Materials and methods

Bacterial strains, plasmids and culture conditions

The bacterial strains and plasmids used in this study are listed in Table 1, whereas oligonucleotide primers are mentioned in Supplementary Material (Online Resource 1). *E. coli* strains were routinely grown in Luria-Bertani (LB) medium at 37 °C (Sambrook et al. 1989). *R. leguminosarum* strains were cultured in 79CA medium with 1 % glycerol as a carbon source (Kowalczyk and Lorkiewicz 1979) and in M1 minimal medium (Sambrook et al. 1989) containing additionally 2 ml l⁻¹ vitamin stock solution (Brown and Dilworth 1995) at 28 °C. When required, antibiotics were used at the following final concentrations: rifampicin, 40 µg ml⁻¹; kanamycin, 40 µg ml⁻¹; tetracycline, 10 µg ml⁻¹; ampicillin, 100 µg ml⁻¹; and spectinomycin, 40 µg ml⁻¹.

DNA methods and sequence analysis

Standard techniques were used for genomic DNA and plasmid isolation, restriction enzyme digestion, cloning,

transformation, hybridization, and sequencing (Sambrook et al. 1989). For PCR reaction, REDTaq Ready PCR Reaction Mix (Sigma) was used. In order to obtain the nucleotide sequence of the Pss cluster, genomic DNA of Rt24.2 and primer walking techniques were used. The resulting sequence of 28 613 bp was deposited in GenBank under the accession no. KP067320. PCR products and plasmid constructs were sequenced using the BigDye terminator cycle sequencing kit (Applied Biosystems) and the ABI Prism 310 sequencer. Database searches were done with the FASTA and BLAST programs available from the National Center for Biotechnology Information (Bethesda, MD, USA) and the European Bioinformatics Institute (Hinxton, UK). Promoter prediction in the *pss* regulatory regions was done using the BDGP Neural Network Promoter Prediction (<http://www.fruitfly.org>). The probability of occurrence of promoter sequences was determined by the use of Promoter prediction (P) score, which can have values from 0 to 1 (a value close to 1 means very high probability, whereas that below 0.7 suggests a lack of promoter sequences). Subsequently, the searches for identification of motifs in the *pss* upstream regions, for which P values were higher than 0.7, were performed with Malign and Fuzznuc programs using *S. meliloti* CTTGAC-N₁₇₋₁₈-CTATAT and *E. coli* TTGACA-N₁₇₋₁₈-TATAAT promoter consensus (MacLellan et al. 2006) as query sequences (<http://www.genebee.msu.su/services/malign>, <http://emboss.ch.embnet.org/Pise>). RosR-box consensus TGAAAT CTAGGGGTAGATTTC A was used as a query sequence for searching of RosR motifs in the *pss* upstream regions (<http://www.genebee.msu.su/services/malign>). Amino acid sequence analyses were performed using the BLASTP program (<http://blast.ncbi.nlm.nih.gov/>) and the CAZy database (www.cazy.org).

Construction of plasmids bearing transcriptional *pss-lacZ* fusions

To construct plasmids containing regulatory regions of the *pss* genes from the Rt24.2 Pss cluster, a broad-host-range plasmid pMP220 carrying a promoterless *lacZ* gene was used. Based on the sequence of this region, primers complementary to promoters of the *pss* genes were designed and used in PCR reactions with the genomic DNA of this strain (Supplementary Material, Online Resource 1). A set of promoter fragments was generated by PCR amplification using primer pairs:

Table 1 Bacterial strains and plasmids used in this study

Strains and plasmids	Relevant characteristics	Sources or reference
<i>R. leguminosarum</i> bv. <i>trifolii</i>		
Rt24.2	Wild type, Rif ^r , Nx ^r	Janczarek et al. 2009
Rt2472	Rt24.2 with a mutation in the <i>rosR</i> gene	Janczarek et al. 2009
Rt24.2(pBR1)	Rt24.2 derivative carrying multiple <i>rosR</i> copies on pBBR1MCS-2 vector	Janczarek et al. 2009
Rt770	Rt24.2 derivative with a mutation in <i>pssS</i> , Sp ^r	This work
<i>E. coli</i>		
DH5 α	<i>supE44</i> Δ <i>lacU169</i> (ϕ 80 <i>lacZ</i> Δ M15) <i>hsdR17</i> <i>recA1</i> <i>endA1</i> <i>gyrA96</i> <i>thi-1</i> <i>relA1</i>	Sambrook et al. 1989
S17-1	294, <i>thi</i> , RP4-2-Tc::Mu-Km::Tn7	Simon et al. 1983
mTn5SS <i>gusA40</i>	miniTn5 interposon with promoterless <i>gusA</i> gene, Sp ^r	Wilson et al. 1995
Plasmids		
pUC19	Cloning and sequencing vector, Ap ^r	Sambrook et al. 1989
pMP220	IncP, <i>mob</i> , promoterless <i>lacZ</i> , Tc ^r	Spaink et al. 1987
pBBR1MCS-2	<i>mob</i> , <i>lacZ</i> α , Km ^r cloning vector (in this study named as pBK1)	Kovach et al. 1995
pBR1	pBBR1MCS-2 with 1.1-kb <i>EcoRI</i> - <i>Bam</i> HI fragment containing the <i>rosR</i> gene	Janczarek et al. 2009
pUC-mgl1	pUC19 with 0.65-kb <i>Bam</i> HI- <i>Sph</i> I fragment of the <i>mgl2</i> promoter region	This work
pUC-V2	pUC19 with 0.9-kb <i>Kpn</i> I- <i>Xba</i> I fragment of the <i>pssV</i> promoter region	This work
pUC-W3	pUC19 with 0.74-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssW</i> promoter region	This work
pUC-M4	pUC19 with 0.65-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssM</i> promoter region	This work
pUC-K5	pUC19 with 0.48-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssK</i> promoter region	This work
pUC-L13	pUC19 carrying the 0.52-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssL</i> promoter region	This work
pUC-I6	pUC19 with 0.9-kb <i>Eco</i> RI- <i>Sph</i> I fragment of the <i>pssI</i> promoter region	This work
pUC-F7	pUC19 with 0.65-kb <i>Bam</i> HI- <i>Pst</i> II fragment of the <i>pssF</i> promoter region	This work
pUC-C8	pUC19 with 0.55-kb <i>Eco</i> RI- <i>Sph</i> I fragment of the <i>pssC</i> promoter region	This work
pUC-D14	pUC19 with 0.35-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssD</i> promoter region	This work
pUC-T9	pUC19 with 0.8-kb <i>Pst</i> II- <i>Bam</i> HI fragment of the <i>pssT</i> promoter region	This work
pUC-N10	pUC19 with 0.75-kb <i>Bam</i> HI- <i>Pst</i> II fragment of the <i>pssN</i> promoter region	This work
pUC-O11	pUC19 with 0.65-kb <i>Bam</i> HI- <i>Pst</i> II fragment of the <i>pssO</i> promoter region	This work
pUC-P12	pUC19 with 0.85-kb <i>Eco</i> RI- <i>Xba</i> I fragment of the <i>pssP</i> promoter region	This work
pUC-R13	pUC19 with 0.64-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssR</i> promoter region	This work
pMgl16	pMP220 carrying the 0.65-kb <i>Bgl</i> II- <i>Sph</i> I fragment of the <i>mgl2</i> promoter region	This work
pV90	pMP220 carrying the 0.9-kb <i>Kpn</i> I- <i>Xba</i> I fragment of the <i>pssV</i> promoter region	This work
pW74	pMP220 carrying the 0.74-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssW</i> promoter region	This work
pM65	pMP220 carrying the 0.65-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssM</i> promoter region	This work
pK48	pMP220 carrying the 0.48-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssK</i> promoter region	This work
pL52	pMP220 carrying the 0.52-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssL</i> promoter region	This work
pI90	pMP220 carrying the 0.9-kb <i>Eco</i> RI- <i>Sph</i> I fragment of the <i>pssI</i> promoter region	This work
pF65	pMP220 carrying the 0.65-kb <i>Bgl</i> II- <i>Pst</i> II fragment of the <i>pssF</i> promoter region	This work
pC55	pMP220 carrying the 0.55-kb <i>Eco</i> RI- <i>Sph</i> I fragment of the <i>pssC</i> promoter region	This work
pT80	pMP220 carrying the 0.8-kb <i>Bgl</i> II- <i>Pst</i> II fragment of the <i>pssT</i> promoter region	This work
pD35	pMP220 carrying the 0.35-kb <i>Eco</i> RI- <i>Pst</i> II fragment of the <i>pssD</i> promoter region	This work
pN76	pMP220 carrying the 0.75-kb <i>Bgl</i> II- <i>Pst</i> II fragment of the <i>pssN</i> promoter region	This work
pO66	pMP220 carrying the 0.65-kb <i>Bgl</i> II- <i>Pst</i> II fragment of the <i>pssO</i> promoter region	This work
pP85	pMP220 carrying the 0.85-kb <i>Eco</i> RI- <i>Xba</i> I fragment of the <i>pssP</i> promoter region	This work

Table 1 (continued)

Strains and plasmids	Relevant characteristics	Sources or reference
pR64	pMP220 carrying the 0.64-kb <i>EcoRI-PstI</i> fragment of the <i>pssR</i> promoter region	This work
pWSR2	pBBR1MCS-2 with the 3.46-kb <i>BamHI-SacI</i> fragment containing <i>pssWSR</i>	This work
pWS4	pBBR1MCS-2 with the 2.9-kb <i>BamHI-SacI</i> fragment containing <i>pssWS</i>	This work
pW5	pBBR1MCS-2 with the 1.7-kb <i>BamHI-SacI</i> fragment containing <i>pssW</i>	This work

Nx^r, nalidixic acid resistance, Rif^r, rifampicin resistance, Tc^r, tetracycline resistance, Ap^r, ampicillin resistance, Sp^r, spectinomycin resistance, Km^r, kanamycin resistance

mgl2-27 and mgl2-28 primers for the pMgl16 plasmid fusion, pssV-69 and pssV-70 for pV90, pssW-29 and pssW-30 for pW74, pssM-33 and pssM-34 for pM65, pssL-101 and pssL-102 for pL52, pssK-31 and pssK-32 for pK48, pssI-35 and pssI-36 for pI90, pssF-37 and pssF-38 for pH65, pssC-39 and pssC-40 for pC55, pssD-41 and pssD-79 for pD35, pssP-42 and pssP-89 for pP85, pssO-43 and pssO-44 for pO66, pssN-45 and pssN-46 for pN76, pssT-47 and pssT-48 for pT80, and pssR-EcoI and pssR-PstI for pR64. The PCR products obtained were digested with appropriate restriction enzymes, cloned into the corresponding sites in the pUC19 vector, and verified by sequencing. Then, plasmid-derived fragments were re-cloned between the respective sites of the pMP220 vector, yielding plasmids with the *pss* upstream regions fused to the promoterless *lacZ* gene. These plasmids were introduced into *E. coli* S17-1 by transformation and subsequently into Rt24.2 derivatives by biparental conjugation (Sambrook et al. 1989).

β-Galactosidase assay

R. leguminosarum derivatives harboring the *pss-lacZ* fusions were grown for 24 h in 79CA medium supplemented with tetracycline and, where necessary, kanamycin. The assay of β-galactosidase activity was carried out according to the protocol described by Miller (Miller 1972). The reported values are given in Miller units and are averages of at least five independent experiments.

Mutagenesis of the *pssS* gene

In order to obtain Rt24.2 derivatives defective in EPS synthesis, random mutagenesis of this strain was performed using the S17-1 strain containing the mTn5SS*gusA40* transposon with a promoterless *gusA* gene (Wilson et al. 1995). As a result of the biparental conjugation experiments, in total 2000 strains carrying

mutations were obtained, among them five clones had a non-mucoid phenotype. The insertion site of the mTn5SS*gusA40* transposon in the genome of the Rt24.2 derivatives was established by restriction, hybridization, and sequencing analyses. Among these, only the one mutant strain, named Rt770 proved to have the transposon insertion located within the *Pss* region, in the *pssS* gene.

Construction of plasmids for complementation of a *pssS* mutation

To construct a set of plasmids containing different fragments of the *pssWSR* region, the pBBR1MCS-2 vector and amplicons obtained in PCR reactions with primers complementary to this region were used (Supplementary Material, Online Resource 1). Primers W-FW3 and W-RW5 were used for construction of a pW5 plasmid, primers W-FW3 and S-RW4 for construction of pWS4, and primers W-FW3 and R-RW3 for construction of pWSR2. The PCR products obtained were digested with *BamHI* and *SacI* enzymes, cloned into the corresponding sites in the pBBR1MCS-2 vector, and inserts of the resulting plasmids were verified by sequencing. Then, the pW5, pWS4, and pWSR2 plasmids were introduced into *E. coli* S17-1 by transformation and subsequently into Rt770 by biparental conjugation.

EPS isolation and quantification

For EPS isolation, 10-ml cultures of rhizobial strains were grown in 79CA medium supplemented with 1 % glycerol for 2 days at 28 °C. Then, the cultures were centrifuged for 20 min at 14,000 rpm. EPS was precipitated from obtained supernatants with 4 vol. of 96 % ethanol, collected by centrifugation, dissolved in 2 ml of water and analyzed for carbohydrates (Loewus 1952). The total sugar content was calculated as glucose equivalents.

Motility assay

The motility of the rhizobial strains was established using 0.3 % 79CA medium. 5- μ l aliquots of bacterial suspensions ($OD_{600}=0.4$) of these strains prepared in sterile water were stabbed into agar. Then, the plates were incubated at 28 °C for 48 h, and the migration distance of bacteria from the injection site was measured. The assay was repeated three times with three repetition for each strain analyzed.

Sensitivity assay to stress factors

In order to determine the sensitivity of the Rt770, Rt24.2 and Rt770(pWS4) strains to several stress factors (sodium dodecyl sulfate, sodium deoxycholate, and ethanol), the minimal inhibitory concentration of the individual stressor was determined. In this purpose, bacterial suspensions of these strains of $OD_{600}=0.2$ were prepared into sterile water and 10- μ l aliquots were placed on the plates with 79CA medium and defined concentrations of the tested compound (SDS - concentrations from 0.05 to 1 % w/v, DOC – from 0.05 to 1 % w/v, ethanol – from 0.05 to 5 % v/v). The bacterial growth on the individual media was determined after 48-h incubation at 28 °C.

Plant experiments

Seeds of red clover (*Trifolium pratense* cv. Diana) were surface sterilized as described previously (Jaszek et al. 2014). Next, the seeds were placed on plates with Fåhræus agar (Vincent 1970), incubated for 2–3 days at 22 °C, and seedlings were subsequently placed onto Fåhræus slants. After 4 days, the seedlings were inoculated with bacterial suspensions with $OD_{600}=0.2$ (100- μ l aliquot per plant). The plants were grown for 28 days under natural light supplemented with artificial light (14 h at 24 °C and 10 h at 18 °C) in a greenhouse. Nodules appearing on the roots were counted after each 7 days. 4-week plants were harvested, and their wet shoot masses were estimated. The experiment was repeated three times using 20 plants for each treatment.

Nodule analysis using light and electron microscopy

To establish nodule colonization by the Rt770 mutant, the enzymatic activity of β -glucuronidase encoded by *gusA* in the mTn5SS*gusA*40 transposon was used. In addition, Rt24.2 tagged with the pJBA21Tc plasmid

carrying *gusA* was used as a control (Janczarek and Rachwał 2013). Clover seedlings were inoculated with these strains and grown up to 3 weeks. Next, the nodules were stained using 50 mM sodium phosphate buffer (pH 7.2) containing 50 μ g ml⁻¹ of 5-bromo-4-chloro-3-indolyl- β -D-glucuronide as described previously (Janczarek and Rachwał 2013) and analyzed under a Nikon light microscope (OPTIPHOT2). In order to characterize in detail the structures of the nodules induced by the Rt24.2 strain and the Rt770 mutant, plant material was prepared for the electron microscopy analysis as described earlier (Król et al. 1998).

Results

The sequence analysis of the Pss region in *R. leguminosarum* bv. *trifolii* 24.2

Rhizobia secrete large amounts of EPS into the environment (D’Haeze et al. 2004; Quelas et al. 2006; Janczarek et al. 2015a). Although some data concerning genetic characteristics of a few genes involved in EPS synthesis in *R. leguminosarum* have been published previously, no comprehensive studies on the transcriptional activity of genes grouped in the EPS synthesis region have been performed so far. Therefore, we decided to establish the genetic organization of this cluster in *R. leguminosarum* bv. *trifolii* strain 24.2 and transcriptional activity of the individual genes present in it. At first, sequencing of the Rt24.2 Pss cluster was performed and the 28.613-kb sequence obtained was deposited in the GenBank database under Accession No. KP067320. Based on the nucleotide sequence analysis, 23 open reading frames (ORFs) were identified in this region, which showed significant identity to genes from other rhizobial species involved in biosynthesis, export, and modification of extracellular polysaccharides. Comparative sequence analysis of this cluster with nucleotide sequences of *R. leguminosarum* bvs. *trifolii* TA1 and *viciae* 3841 strains as well as closely related species *Rhizobium etli* CFN42 indicated that this DNA region showed high synteny (Fig. 1). The *pss* genes of the Rt24.2 cluster showed the same genetic organization as that found in the RtTA1, Rv3841, and ReCFN42 bacteria. Only its fragment encompassing the *plyA*, *rapA*, and *orf1* genes proved to be more divergent, and this difference was especially visible in the case of the RtTA1 strain. The entire Rt24.2 Pss cluster showed the highest nucleotide

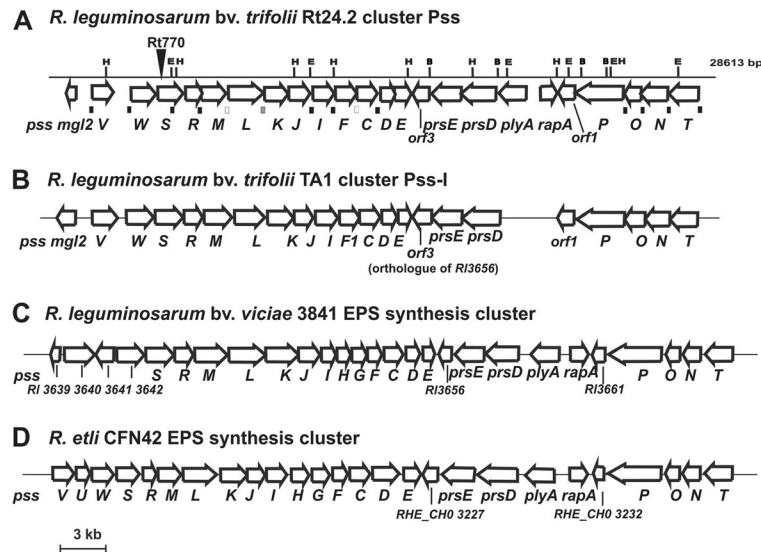


Fig. 1 Genetic organization of exopolysaccharide synthesis clusters in different rhizobia. **a** Physical and genetic map of the Pss region of *R. leguminosarum* bv. *trifolii* 24.2. Arrows below the map indicate the direction of transcription. Promoter regions of the *pss* genes cloned into the pMP220 vector and present in the particular *pss-lacZ* transcriptional fusions are marked as rectangles: black – DNA regions characterized by very high probability of the promoter occurrence (*P* values) higher than 0.9, grey – regions with the *P* values higher than 0.8, white – regions with

the *P* values higher than 0.7 (<http://www.fruitfly.org>). Selected restriction sites are marked: E- *EcoRI*, H- *HindIII*, B- *BamHI*. Location of the mTn5SSgusA40 insertion in the Pss region of *R. leguminosarum* bv. *trifolii* mutant Rt770 is marked by dark grey vertical arrow. **b** Genetic map of the Pss-I region of *R. leguminosarum* bv. *trifolii* TA1, **c** Genetic map of the *R. leguminosarum* bv. *viciae* 3841 EPS synthesis cluster, **d** Genetic map of the *R. etli* CFN42 EPS synthesis cluster

sequence identity to the homologous region of RtTA1 (95 % identity), and a slightly lower level to the sequences of Rv3841 (91 %) and ReCFN42 (83 %) strains, confirming that these genetic regions are highly conserved among these rhizobial strains. Depending on putative functions in EPS biosynthesis, the *pss* genes of the Rt24.2 Pss cluster were divided into 3 groups. Genes involved in the subunit synthesis were classified into the first group: *pssJ* encoding galactosyltransferase, *pssS*, *pssI*, and *pssF* coding for glycosyltransferases, and *pssC*, *pssD*, and *pssE* genes coding for glucuronosyltransferases. The second group included genes involved in addition of non-sugar residues to the EPS subunits: *pssR* for acetyltransferase and two *pssM* and *pssK* genes for ketal pyruvate transferases. The third group encompassed the genes involved in EPS polymerization and export: *pssP* encoding an EPS polymerization protein, *pssO* encoding an outer membrane protein, and the *pssN*, *pssT* and *pssL* genes encoding Wza-type, Wzy-type and Wzx-type polysaccharide export proteins, respectively. Moreover, other genes not directly involved in EPS biosynthesis were identified in the Rt24.2 Pss cluster. These were: *pssW* encoding a glycosyl hydrolase probably engaged in EPS processing, the *prsD* and *prsE* genes encoding

components of the type-I protein secretion system and three genes, named *plyA*, *rapA*, and *orf3*, which encode proteins that are substrates for this transport system.

In conclusion, the Rt24.2 Pss cluster shows very similar genetic organization with those of other *R. leguminosarum* strains, although some minor differences were found.

Transcriptional activity of the *pss* genes located in the Rt24.2 Pss region

In order to identify putative promoters of the *pss* genes present in the Rt24.2 Pss cluster, *in silico* sequence analysis of upstream regions of these genes was performed using both *S. meliloti* (5'-CTTGAC-N₁₇₋₁₈-CTATAT-3') and canonical *E. coli* (5'-TTGACA-N₁₇-TATAAT-3') promoter consensus motifs recognized by RNA polymerase with ⁷⁰. It was established that a great majority of the analyzed genes possessed motifs in their regulatory regions of a high probability (*P*) of occurrence of promoter sequences (*P* score higher than 0.9 for *mgl2*, *pssV*, *pssW*, *pssR*, *pssM*, *pssI*, *pssF*, *pssP*, *pssO*, *pssN*, *pssD* and *pssT*). The values between 0.8 and 0.7 for *pssL*, *pssK*, *pssD*, and *pssC* suggested that these

genes had motifs of the significantly lower probability of presence of promoter sequences. In the case of the upstream regions of the *pssS*, *pssJ*, and *pssE* genes, no motifs of the predicted promoter activity were identified. Next, the regulatory regions of these genes, for which the determined P scores were higher than 0.7, were amplified in PCR reactions and cloned upstream of a promoterless *lacZ* gene into the pMP220 vector. As a result, a set of 15 *pss-lacZ* transcriptional fusions containing the promoter regions of the different genes from the Rt24.2 Pss cluster were constructed. These plasmids were introduced into the wild-type Rt24.2 strain and β -galactosidase activity was assayed (Table 2).

In general, for almost all of the regulatory *pss* regions chosen for cloning, the presence of functional promoters in their sequences was confirmed. Moreover, high diversity concerning the levels of the promoter activity was observed among these genes. The highest transcription was observed for *pssN*, *pssO*, *pssR*, *mgl2*, and *pssT*, indicating that the upstream regions of these genes

contain strong promoters (Table 2). Lower values of β -galactosidase activities were found for the *pssI-lacZ*, *pssP-lacZ*, *pssV-lacZ*, *pssF-lacZ*, and *pssW-lacZ* fusions (a range from 1106 to 661 Miller units). The lowest transcriptional activity was detected for the *pssM*, *pssC*, *pssK*, and *pssD* genes (from 325 to 157 Miller units), confirming the presence of weak promoters in their sequences. In the case of the *pssL-lacZ* fusion, the level of β -galactosidase activity was nearly the same as that determined for the control plasmid pMP220, indicating that no promoter sequence was present in the upstream region of this gene. Based on all these results, it was confirmed that the upstream regions of the *pssV*, *pssW*, *pssR*, *pssM*, *pssK*, *pssI*, *pssF*, *pssC*, *pssD*, *pssT*, *pssN*, *pssO*, *pssP*, and *mgl2* genes contain functional promoters, although of different strengths of action. In addition, we conclude that the *pssV*, *pssI*, *pssF*, *pssC*, *pssR*, *pssT*, *pssN*, *pssO*, *pssP*, and *mgl2* genes constitute single transcription units, whereas the transcription of other *pss* genes is linked (*pssWS*, *pssML*, *pssKJ*, and *pssDE* operons).

Table 2 Transcriptional activity of the *pss-lacZ* fusions assayed in the *R. leguminosarum* bv. *trifolii* wild-type strain 24.2 and its derivatives Rt2472*rosR*, Rt24.2(pBR1) and Rt24.2(pBK1)

β -galactosidase activity (Miller units)							
Plasmid	<i>pss-lacZ</i> fusion	Rt24.2 (wt)	Rt2472 (<i>rosR</i> ⁻)	Ratio Rt2472/Rt24.2	Rt24.2 (pBK1)	Rt24.2 (pBR1)	Ratio Rt24.2(pBR1)/Rt24.2(pBK1)
pMgl6	<i>mgl2-lacZ</i>	1877±121	2083±168	1.11	1853±125	1798±174	0.97
pV90	<i>pssV-lacZ</i>	1056±87	1124±109	1.06	1009±90	1087±113	1.08
pW74	<i>pssW-lacZ</i>	661±87	821±103	1.24	642±78	568±83	0.88
pR64	<i>pssR-lacZ</i>	4014±1012	3813±889	0.95	3902±891	4214±923	1.08
pM65	<i>pssM-lacZ</i>	325±29	367±38	1.13	291±37	304±28	1.04
pK48	<i>pssK-lacZ</i>	247±19	293±33	1.19	234±22	252±22	1.07
pL56	<i>pssL-lacZ</i>	54±8	59±10	1.08	51±9	57±9	1.11
pI90	<i>pssI-lacZ</i>	1106±91	1198±121	1.08	1121±103	1041±121	0.93
pF65	<i>pssF-lacZ</i>	794±51	1664±143	2.09*	781±66	552±57	0.71*
pC55	<i>pssC-lacZ</i>	263±24	248±28	0.94	251±30	255±27	1.01
pD35	<i>pssD-lacZ</i>	157±19	164±23	1.05	144±17	148±21	1.03
pT80	<i>pssT-lacZ</i>	1386±111	1538±149	1.11	1217±141	1289±119	1.06
pN76	<i>pssN-lacZ</i>	12,311±971	18,712±1397	1.52*	12,191±902	9145±693	0.75*
pO66	<i>pssO-lacZ</i>	7294±514	12,837±989	1.76*	7366±634	5250±387	0.71*
pP85	<i>pssP-lacZ</i>	1061±102	2005±207	1.89*	1019±98	711±92	0.69*
pMP220	—	39±7	48±8	—	39±7	42±6	—

The bacteria were grown in 79CA medium supplemented with 1 % glycerol for 24 h. Data shown are the mean±SD

* Asterisks indicate a statistically significant differences between the wild-type Rt24.2 and the mutant strain Rt2472, and between the Rt24.2(pBR1) and the Rt24.2(pBK1), respectively (*P* value <0.05; Student's *t* test)

Promoter sequences of the *pss* genes

Up to now, the ability to recognize and predict non-⁵⁴ promoters in alphaproteobacteria has not been well developed. Among rhizobial species, the data concerning this issue were established in detail only for *S. meliloti*, in which consensus 5'-CTTGAC-N₁₇₋₁₈-CTATAT-3' based on 25 experimentally verified promoter sequences was established (MacLellan et al. 2006). The structure of this consensus shares significant similarity with the *E. coli*⁷⁰ consensus (5'-TTGACA-N₁₇₋₁₈-TATAAT-3'), but is skewed by 1 nucleotide to the left.

In this study, a comparative analysis of the regulatory sequences of the *pss* genes was performed using both *E. coli* and *S. meliloti*⁷⁰ promoter consensus. These data are presented in Table 3. In general, a majority of the promoter sequences of the *R. leguminosarum pss* genes shared significant similarity with the *S. meliloti*⁷⁰ consensus (*pssV*, *pssW*, *pssF*, *pssC*, *pssD*, *pssT*, *pssO*, *pssR*, *pssN*, and *pssP*). Moreover, it was established that the -35 motifs of the *pss* promoters had higher identity than the -10 motifs with the corresponding hexamers of the *S. meliloti* consensus. To find reasons for the extremely high promoter activity of *pssN* and *pssO*, we additionally searched the upstream regions of these

genes for identification of other regulatory motifs affecting the level of transcription such as an upstream promoter element (UP) and extended -10 element. It is well known that these -10 elements significantly enhance the promoter strength (Ross et al. 1998; Estrem et al. 1999). The 30-bp-long A/T-rich UP sequence is located upstream of the -35 hexamer, whereas the 3-bp TGN motif upstream of the -10 hexamer. The A/T-rich sequences were found upstream of the *pssN* -35 motif (5'-ACAA TCAATCTTTTCCGAAATCTTATTAATCCATC CATTAAACTTTAA-3') and the *pssO* -35 motif (5'-AACTTGACCAATTCAACAGAGTCCGAATTGGA AATATAGATAATATCCTTGTCTT-3'). This indicated the significant role of these UP elements in enhancing transcription of these rhizobial genes and explained the highest activity of these promoters among all the studied *pss* promoters.

The role of RosR in regulation of expression of the *pss* genes

Previously, it was established that RosR positively affected EPS production in *R. leguminosarum* (Janczarek and Skorupska 2007; Janczarek et al. 2009). Therefore, we decided to determine whether RosR regulates EPS

Table 3 Predicted -35 and -10 sequences of the *pss* gene promoters

Promoter sequence (5'-3')							
No	Gene	1 nt upstream of the -35 motif	-35 hexamer	17-18 nt	-10 hexamer	1 nt downstream of -10 motif	Distance to translation start (nt)
1	<i>mgl2</i>	T	TCGACA	GATTATTGTGTGTATTA	TGATAT	G	-257
2	<i>pssV</i>	C	TTGATC	TAAAGCGCGTCGCGATC	TTTTAG	A	-227
3	<i>pssW</i>	A	TTGACG	CAGCCGCCACGACATTC	TGACGA	G	-368
4	<i>pssR</i>	C	TTGCCA	AGCGTCACGATCCATGA	AAATCT	G	-484
5	<i>pssM</i>	T	TTGTGA	AGGCCAAGTGGCTCTAC	TATACG	A	-259
6	<i>pssK</i>	G	TCGTCA	CCGGCGGTATCCTCTAT	GTCGGT	T	-223
7	<i>pssI</i>	T	TTCGCA	ATCTTCGCAAGCTGAGG	GTACTC	G	-202
8	<i>pssF</i>	C	TGGATA	CCGGCATCAAGTGTCAC	TGAGGA	T	-285
9	<i>pssC</i>	A	TTGCCG	AAACGACATCGCTTCTG	AAGCTC	C	-116
10	<i>pssD</i>	C	TTGGAA	GAGTTTGACCCCTTATT	ATGTTT	C	-181
11	<i>pssT</i>	A	TTGGCT	TAAACCTATCGCGACAA	TTTTAA	A	-86
12	<i>pssN</i>	G	TTGACT	CTGAAACCCAAAGAAATT	TTATAG	C	-197
13	<i>pssO</i>	C	TTGACA	TCGTAAACTCGGTATCG	TCCACT	G	-136
14	<i>pssP</i>	C	TTCGAG	TTCCACGACCACGACTAC	GATCAT	C	-129
<i>E. coli</i> ^a		—	TTGACA		TATAAT	—	
<i>S. meliloti</i> ^a		C	TTGAC—	C	TATAT	C	

^a *E. coli* and *S. meliloti* consensus sequences

synthesis via modulation of transcription of the *pss* genes present in the *Pss* cluster. In order to verify this hypothesis, the set of the *pss-lacZ* fusions was introduced into the Rt2472*rosR* mutant, the Rt24.2(pBR1) strain containing additional *rosR* copies, and the Rt24.2(pBK1) strain containing the empty pBBR1MCS-2 vector, and β -galactosidase activity was measured (Table 2). In a majority of the studied *pss-lacZ* fusions, similar levels of β -galactosidase activity were observed in the four strains used: Rt24.2, Rt2472, Rt24.2(pBR1) and Rt24.2(pBK1) (which corresponded to values of nearly 1.0 for both Rt2472/Rt24.2 and Rt24.2(pBR1)/Rt24.2(pBK1) ratios). This indicated that the RosR protein did not influence the transcription of these genes. However, significant differences in the transcription levels of the *pssO*, *pssN*, *pssP*, and *pssF* genes were detected, when the β -galactosidase activities for these individual *pss-lacZ* fusions were compared between Rt24.2 and Rt2472, and between Rt24.2(pBR1) and Rt24.2(pBK1) strains, respectively (Table 2). Among these genes, the highest Rt2472/Rt24.2 ratio was found for the *pssF-lacZ* fusion (2.09). For the remaining genes, slightly lower Rt2472/Rt24.2 ratios were established. On the other hand, the Rt24.2(pBR1)/Rt24.2(pBK1) ratios were determined for these fusions to be significantly lower than 1.0, confirming that multiple *rosR* copies in the Rt24.2(pBR1) strain affected expression of these genes more strongly than the one *rosR* copy in the wild-type strain. These data indicated that the RosR protein was engaged in regulation of transcription of the *pssO*, *pssN*, *pssP*, and *pssF* genes.

In previous studies, it was confirmed that RosR recognized a sequence motif, called the RosR-box, located in the upstream regions of the *rosR* and *pssA* genes and affected their transcription (Janczarek and Skorupska 2007; Janczarek and Urbanik-Sypniewska 2013). The RosR-box is a 22-bp-long sequence containing 9-bp inverted repeats separated by a 4-bp spacer (Fig. 2). In order to establish whether the upstream regions of the *pssO*, *pssN*, *pssP*, and *pssF* genes possess RosR-box motifs, the promoter sequences of these genes were analyzed. We found motifs with significant sequence similarity to the RosR consensus in the *pss* upstream regions, which were located downstream of the promoters and upstream of translation start sites (with the exception of *pssN*) (Fig. 2). Among these, the motif identified in the *pssF* upstream region showed the highest sequence identity to the RosR consensus. Also,

	9 nt	9 nt	Distance from the ORF (nt)
<i>rosR</i> consensus	←TGAAATCTAGGGGTAGATTTC→		
RosR-box upstream of the <i>rosR</i> gene	CGGAATCTAGGGGTGGATTTCG		-111
<i>pssF</i>	GTGGATATCGTCTGGAATTCG		-55
<i>pssP</i>	TAAGATCTACGGGCA-ATTCCAG		-29
<i>pssN</i>	TGGCAACGATGTGCAGCTTGCC		+62
<i>pssO</i>	CCGGAACAGTGGACGATACCG		-130

Fig. 2 Comparison of sequence motifs identified in the *pss* genes that show similarity to RosR-box consensus and the binding site in the *rosR* upstream region. Nucleotides conserved to the consensus sequences are underlined. Arrows indicate inverted repeats

motifs found in the regulatory regions of *pssO*, *pssN*, and *pssP* showed significant similarity to the consensus sequence, suggesting that RosR influenced transcription of these genes via the RosR-boxes.

Phenotype of the Rt770 mutant strain and complementation of a *pssS* mutation

In order to establish mutations in the *pss* genes which totally abolish EPS synthesis, random mutagenesis of the Rt24.2 strain was performed using a mTn5SSgusA40 transposon (Wilson et al. 1995). As a result, about 2000 derivatives were obtained, among them only five strains were unable to produce EPS. Based on the PCR reaction, hybridization, and sequencing analyses, it was established that only one strain, named Rt770, had transposon insertion inside the *Pss* cluster (in the *pssS* gene) (Fig. 1). In addition, the presence of the one copy of the mTn5SSgusA40 transposon in the Rt770 genome was confirmed using hybridization. This mutant formed characteristic small nonmucoid colonies on agar plates, which significantly differed from those formed by wild-type bacteria (data not shown). Quantitative analysis indicated that the Rt770 strain did not produce any amounts of EPS (0 ± 0 mg L⁻¹ in comparison with the 1281 ± 157 mg L⁻¹ produced by Rt24.2). Moreover, although the Rt770 strain grew nearly as effectively as the parental strain in both the energy-rich media (79CA and TY), it exhibited approx. 3-fold slower growth in the minimal M1 medium ($OD_{600} = 0.27 \pm 0.03$ for Rt770 and $OD_{600} = 0.84 \pm 0.06$ for Rt24.2 in the 72-h cultures). Furthermore, the Rt770 strain was characterized by significantly reduced motility and increased sensitivity to several stress factors (Table 4). This mutant was more sensitive to such compounds as SDS, DOC, and ethanol,

Table 4 Motility and sensitivity of the wild-type *R. leguminosarum* bv. *trifolii* 24.2 and its mutant strain Rt770 to various stress factors

Strain	Motility assay (mm) ^b	Minimal inhibitory concentration ^a		
		SDS (% w/v)	DOC (% w/v)	Ethanol (% v/v)
Rt24.2 (wt)	20±2.0 ^A	0.025±0.005 ^A	0.10±0.005 ^A	5.0±0.25 ^A
Rt770 (<i>pssS</i>)	6.0±1.0 ^B	0.010±0.005 ^B	0.075±0.005 ^B	3.5±0.25 ^B
Rt770(pWS4)	19±2.0 ^A	0.025±0.005 ^A	0.09±0.005 ^A	4.50±0.50 ^A

^a Given values are averages±SD of three independent experiments with 3 biological repetitions for each strain and treatment

^b Bacterial migration was determined after 2-day incubation at 28 °C by measuring the distance from the injection site into agar. The data in the column followed by different upper case letters are significantly different ($P<0.05$; ANOVA, post hoc Tukey's test)

indicating that a lack of EPS resulted in a decrease of adaptation ability of this bacterium to stress conditions.

The complementation analysis of the mutation in the Rt770 strain performed by us using pBBR1MCS-2 derivatives containing different fragments of the *pssWSR* region indicated that a mucoid phenotype (Exo⁺) of this mutant was recovered after introduction of pWSR2 and pWS4 plasmids, but not by the pW5 plasmid (Fig. 3). The pWS4 plasmid containing the *pssWS* genes without *pssR* complemented the mutation from the Exo⁻ to Exo⁺ phenotype. These data confirm that *pssS* plays a significant role in EPS synthesis and that the mutation in this gene does not have a polar effect on downstream-located *pssR*. The *pssS* gene encodes a 383-aa long protein, which according to homology search data was referred to the glycosyltransferases from GTB-type superfamily, GT family 1 (EC 2.4.1.1-, CAZy database), which form glycosidic bonds with stereochemistry identical to that of the glycosyl donor. The Rt770 strain harboring pWS4 produced a similar amount of EPS (1397±172 mg L⁻¹) as Rt24.2 (1281±157 mg L⁻¹), and showed similar motility and sensitivity to the tested stress factors as the wild-type strain (Table 4).

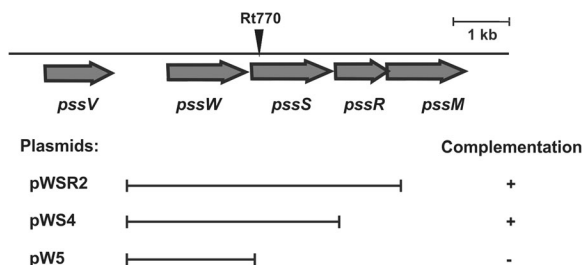


Fig. 3 Complementation analysis of the *pssS* mutation in the Rt770 strain using a set of plasmids harboring different fragments of the *pssWSR* region. + indicates a positive result of complementation (i.e., recovery of the ability of EPS production), whereas – indicates a lack of EPS synthesis

Symbiotic properties of the Rt770 strain

In addition, symbiotic properties of the Rt770 strain in interaction with clover plants were characterized. It was found that this mutant had dramatically reduced infection effectiveness. After 14 days post infection, only 5 % of the clover plants were infected by this bacterium, and less than half of the plants were infected even after 4 weeks (Table 5). Besides the significant delay in nodule formation, also the total number of nodules induced by the *pssS* mutant was essentially lower than that induced by the wild-type strain (4-fold for the 28-day-old plants). Moreover, the nodules elicited by the mutant were small, white, often with the atypical shape and unable to fix nitrogen. This was confirmed by the very low fresh shoot mass of the clover plants, which was 2-fold lower than the mass of the plants infected by the wild-type bacteria, and nearly the same as that of uninfected plants (Table 5).

In order to compare colonization of nodules by Rt770 and Rt24.2, bacteria of these strains tagged with *gusA* encoding β -glucuronidase were used. It was confirmed in this experiment that the wild-type strain colonized the nodules effectively and all zones with the exception of the meristem were occupied by the bacteria (Fig. 4a and b). In contrast, nodule colonization by the Rt770 strain was drastically decreased. A great majority of the nodules were not occupied by the mutant bacteria, which were visible only on the nodule surface (Fig. 4c). In sporadically formed nodules, the bacteria were found inside only single nodule cells (Fig. 4d).

The nodules induced by the wild-type strain Rt24.2 on clover roots exhibited a typical structure with all zones, including a large nitrogen-fixation zone (Fig. 5a). Infection threads had normally formed a thread wall with a large amount of thread matrix

Table 5 Symbiotic properties of *R. leguminosarum* bv. *trifolii* 24.2 and its derivatives

Strain	Infection effectiveness (%) ^a (dpi) ^b				Nodule no. per plant (dpi) ^b				Shoot biomass (mg FW plant ⁻¹)
	7	14	21	28	7	14	21	28	
Rt24.2(wt)	20±4	70±6	100±0	100±0	0.4±0.1	2.5±0.3	5.6±1.1	9.9±2.3	58.7±10.5
Rt770(<i>pssS</i>)	0±0	5±5	35±5	45±10	0±0	0.15±0.05	1.45±0.2	2.5±0.5	29.3±6.7
Rt770(<i>pWS4</i>)	17±3	72±8	100±0	100±0	0.35±0.05	2.1±0.2	5.3±0.5	9.4±0.7	56.2±8.3
Uninoculated clover	–	–	–	–	–	–	–	–	28.9±5.6

Given values are means±SD of three independent experiments with 20 plants for each treatment

^a The values were calculated as a number of plants having nodules on their roots per a number of the plants tested in the experiment

^b days post infection

(Fig. 5b), and mature infected plant cells contained properly differentiated bacteroids (Fig. 5c).

In contrast, 3-week-old nodules induced by the *pssS* mutant were surrounded by cortex built of large, loosely arranged cells and by an endodermis. Semi-thin sectioning of these nodules revealed that a considerable part of central nodule tissue contained uninfected parenchymatous cells with starch grains and only a few cells were infected. The infected cells often formed clusters separated from one another by uninfected parenchyma. The nodules contained the meristem and vascular bundle

connected with the root stele (Fig. 6a). The walls of root epidermal cells, being in contact with the mutant bacteria, were thickened and stained intensely with Azur A and Methylene Blue. The root cortical cells penetrated by infection threads were noticeably enlarged and their walls were also thicker in comparison to other cortex cells (Fig. 6b). The mutant-induced infection threads were wide and often branched. The thread walls were thick, irregularly formed, with knobs and protrusions, which was a consequence of incorporation of additional wall material (Fig. 6c). Another striking morphological feature

Fig. 4 Light microscopy of nodules induced on clover roots (*Trifolium pratense*) by the *R. leguminosarum* bv. *trifolii* wild-type strain 24.2 and the *pssS* mutant Rt770 harboring the *gusA* reporter gene for β -glucuronidase. (**a, b**) Rt24.2 wild-type nodules at 7 and 14 days post infection, respectively; bars=0.15 and 0.4 mm (**c, d**) Rt770 nodules at 6 and 14 days post infection, respectively; bars=0.25 and 0.6 mm. The nodules were stained for GUS activity

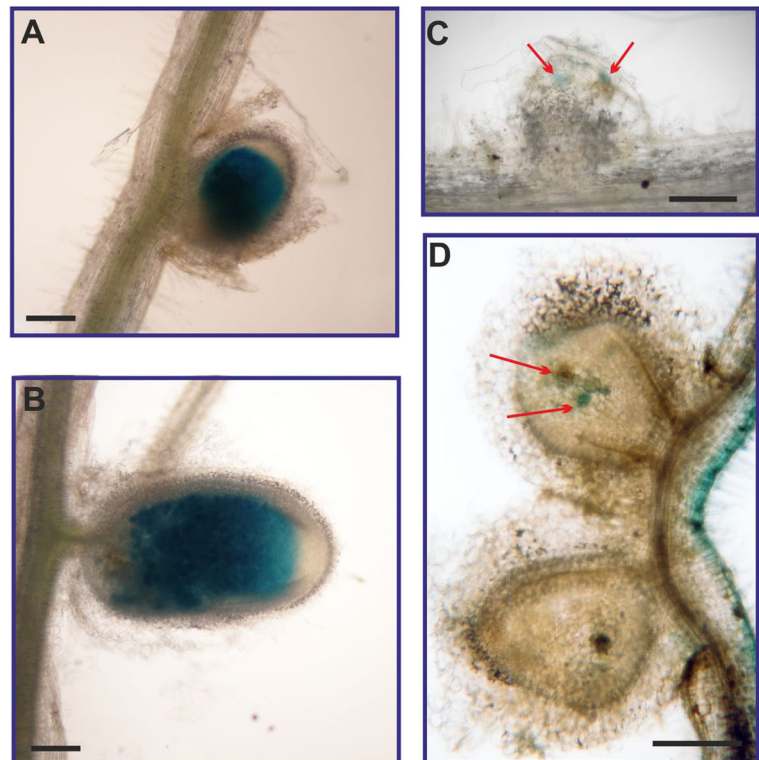
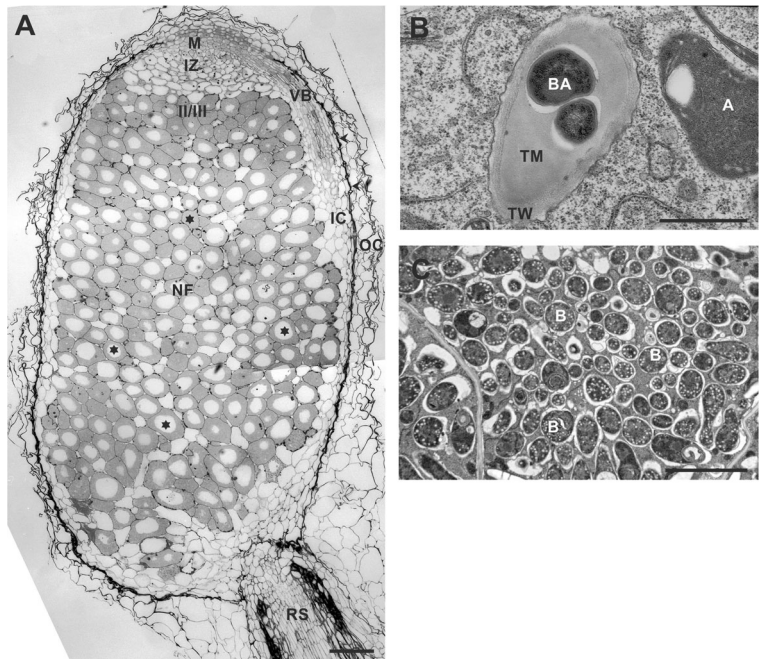


Fig. 5 Wild-type clover root nodule. **a** Semithin section of a 3-week-old nodule induced by the *R. leguminosarum* bv. *trifolii* wild-type strain 24.2: OC - outer cortex; IC - inner cortex; VB - nodule vascular bundle; RS - root stela; M - meristem; IZ - infection zone; II/III - interzone; NF - nitrogen fixation zone; *arrow heads* - nodule endodermis; *asterisks* - infected cells; bar=100 μ m. **b** Ultrastructure of the infection thread: TW - thread wall; TM - thread matrix; BA - bacteria; A - amyloplast; bar=1 μ m. **c** The mature infected plant cell with properly differentiated bacteroids (B); bar=5 μ m



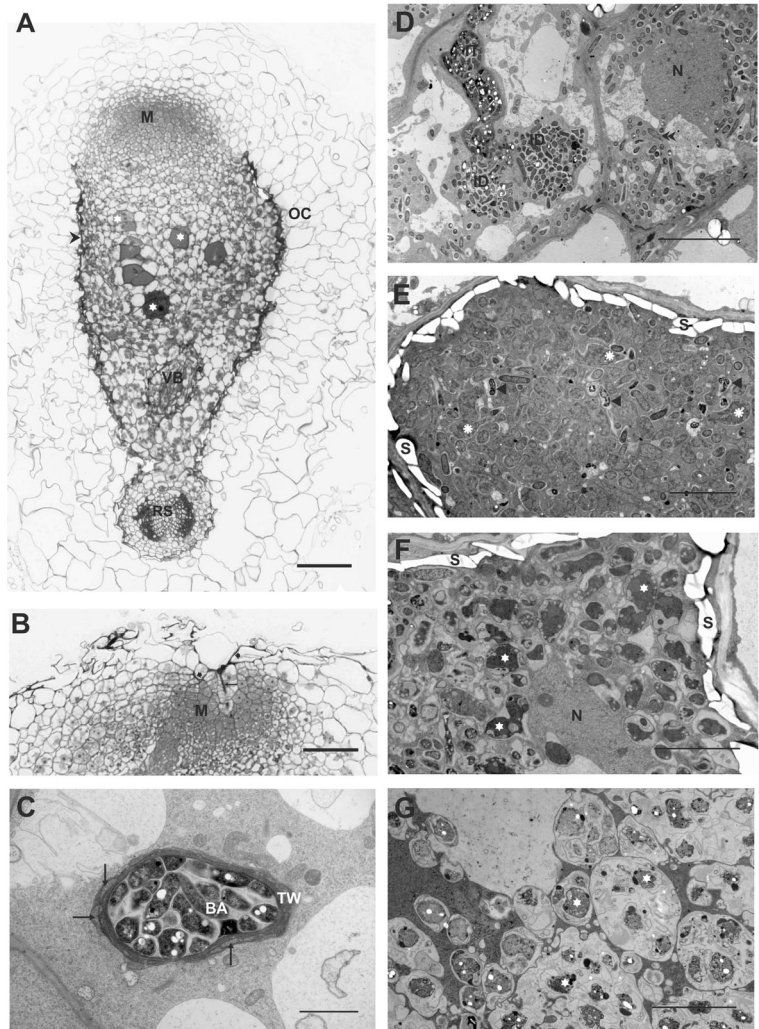
of these infection threads was lack of typical thread matrix, in which rhizobia are usually embedded in the wild-type nodules. The bacterial cells were tightly packed inside these infection threads and showed high variability in their shape and size. The process of plant cell infection was abnormal, being characterized by “explosion-like” mass endocytosis of the bacteria into the plant cell cytoplasm (Fig. 6d). The symbiosomes formed in these cells contained a single bacteroid, and those with more than one bacteroid were observed only occasionally (Fig. 6e). Microscopic analysis revealed that some bacteroids differentiated, while others degenerated precociously. The morphologically pronounced bacteroid differentiation was different from that observed in wild-type nodules. The bacteroids were larger than normal, abnormally swollen and often deformed, and underwent rapid senescence. Their homogenous cytoplasm became electron-dense, marbled or completely dark (black) (Fig. 6f). The bacteroids degenerated inside the peribacteroid membranes or, when the peribacteroid membranes were fused, were aggregated in lytic compartments (Fig. 6g).

In conclusion, all these data indicate that symbiosis of the Rt770 strain with clover was impaired in the early stages of symbiosis, i.e., the infection process. The mutant bacteria infected only sporadic nodule cells, and bacteroids inside these plant cells degenerated early, which in consequence made the nodules unable to fix nitrogen.

Discussion

Rhizobia belong to a very important group of soil bacteria which possess a unique ability of establishing nitrogen-fixing symbiosis with legumes (Broughton et al. 2003). This type of the plant-microbe interaction is essential for biosphere functioning, since it provides nutrients to plants, increases soil fertility and restores deranged/damaged ecosystem (Kennedy et al. 1997; Yanni et al. 1997; Qin et al. 2011; Abreu et al. 2012; Angus et al. 2013). The process of biological nitrogen fixation is carried out by microsymbionts in specialized structures formed on legume roots, called nodules. However, rhizobia must survive often a long period of time in the soil as free-living organisms until they meet their symbiotic partner. During this period, they are exposed to the action of several environmental factors (e.g., drought, pH, temperature, salinity, nutrient limitation, heavy metals, oxidative stress) (Frey and Blum 1994; R s nen et al. 2004; Soussou et al. 2013; Talbi et al. 2013; Suzuki et al. 2014; Cazenave et al. 2014; Pini et al. 2014; Burchill et al. 2014; Barthelemy-Delaux et al. 2014; Jaszek et al. 2014). Extracellular polysaccharide secreted in large amounts by rhizobial cells plays a significant protective role in their adaptation to stress conditions. Several environmental factors such as the type of carbon source, nitrogen and phosphorus

Fig. 6 Semithin section of a 3-week-old clover root nodule induced by the *R. leguminosarum* bv. *trifolii* mutant Rt770 (**a**). M - meristem; OC - outer cortex; RS - root stela; VB - nodule vascular bundle; *white asterisks* - infected cells; *arrow head* - nodule endodermis; bar=100 μ m. **b**) The cortical region of the nodule induced by the mutant Rt770: M - meristem; *long arrow* - infection thread; *short arrows* - thickened cell walls of root epidermis and cortical cells; bar=100 μ m. **c**) Ultrastructure of the infection thread: BA - bacteria; TW - thread wall; *arrows* - material deposited between osmiophilic layers of the thread wall; bar=2 μ m. **d**) “Explosion-like” mass endocytosis of bacteria into the plant cytoplasm: ID - infection droplet; IT - infection thread; N - nucleus; *double arrow heads* - young bacteroids; bar=10 μ m. **e**) Mature infected cell: S - starch granules; rosette - abnormally differentiated bacteroids; *triangle* - degenerated bacteroids; bar=5 μ m. **f**) The infected cell with adult degrading bacteroids (*stars*); N - nucleus; S - starch granules, bar=5 μ m. **g**) Lytic compartments with degrading bacteroids (*stars*); bar=5 μ m



availability, salinity, pH, and temperature, affect the level of EPS produced by rhizobia (Quelas et al. 2006; Janczarek and Skorupska 2009, 2011; Huang et al. 2012; Janczarek et al. 2015a). The synthesis of EPS is a multi-step process, which involves a coordinated action of many enzymatic proteins. In *R. leguminosarum*, a great majority of genes encoding these proteins are located in a large chromosomal cluster designated Pss (Polysaccharide synthesis). In this study, the Pss region of *R. leguminosarum* bv. *trifolii* strain 24.2 was sequenced and genetically characterized. It displayed high synteny with the homologous regions of other strains from *R. leguminosarum* species (Rv3841, RtTA1) and closely related bacterium *R. etli* CFN42 (Fig. 1) (Young et al. 2006; Król et al. 2007; González et al. 2006). In general, the organization and the genetic content of these regions are very similar, and the sequence identity

is very high. The detailed sequence analysis of the Rt24.2 Pss cluster resulted in identification of promoters located upstream of the *pss* genes (Table 3), which indicated significant similarity with the *S. meliloti* or canonical *E. coli* ⁷⁰ promoters (MacLellan et al. 2006; Schlüter et al. 2013). Based on levels of the transcriptional activity of the *pss-lacZ* fusions, we conclude that the *pss* genes encoding enzymatic proteins involved in the synthesis and modification of EPS are transcribed at low levels (e.g., *pssDE*, *pssWS*, *pssC*, and *pssKJ*), whereas those encoding proteins engaged in polymerization and export of EPS have stronger promoters (*pssN*, *pssO*, *pssT*, and *pssP*). This was especially visible for the *pssN* and *pssO* genes, whose upstream regions contained the additional regulatory element UP (Estrem et al. 1999; Browning and Busby 2004). Up to now, UP elements increasing promoter activity were described in

only a small number of bacterial promoters, among them the *rosR* promoter (Aiyar et al. 1998; Ross et al. 1998; Janczarek and Skorupska 2009).

Our previous studies indicated that RosR plays an essential role in positive regulation of EPS synthesis in *R. leguminosarum*, since a *rosR* mutation significantly decreases the production of this polysaccharide. RosR is a *Rhizobium*-specific transcriptional regulator, which binds to a sequence motif, called the RosR-box located in the upstream regions of the regulated genes. Homologs of the Rt24.2 RosR were found in other closely related bacteria such as *R. etli* (Bittinger et al. 1997), *A. radiobacter* (D'Souza-Ault et al. 1993), *A. tumefaciens* (Chou et al. 1998; Malgieri et al. 2007), and *S. meliloti* (Rüberg et al. 1999). Therefore, we decided to establish whether expression of some *pss* genes located in the Rt24.2 Pss cluster is regulated by RosR. For this purpose, transcriptional activities of the individual *pss-lacZ* fusions were compared between the wild-type and the Rt2472*rosR* mutant, and between Rt24.2(pBR1) containing multiple *rosR* copies and Rt24.2(pBK1) containing the empty vector pBBR1MCS-2, respectively. Statistically significant differences in expression levels were found for the *pssF*, *pssP*, *pssO*, and *pssN* genes (Rt2472/Rt24.2 and Rt24.2(pBR1)/Rt24.2(pBK1) ratios) (Table 2). In the upstream regions of these genes, motifs resembling the RosR-box were found, suggesting that these genes belong to the RosR regulon (Fig. 2). Similarly in *S. meliloti*, MucR, which is a RosR homolog, affected the synthesis of both exopolysaccharides: succinoglycan and galactoglucan via repression of some genes (*wgaA*, *wgdA*, *wgeA*, *wggR*, *exoH*, and *exoX*) and activation of other genes (*exoYFQ* and *exoK*) (Bertram-Drogatz et al. 1998; Quester and Becker 2004). Also in *A. tumefaciens*, the RosR homolog (Ros) negatively regulated expression of *virC* and *virD* genes, which positively affected EPS production in this bacterium (Chou et al. 1998).

The production of EPS is very important not only for free-living rhizobia, but also for establishment of effective symbiosis with many legumes (e.g., clover, alfalfa, vetch, and peas) (Ivashina et al. 1994; Cheng and Walker 1998; Janczarek and Rachwał 2013). In this study, we have characterized a strain Rt770 having a mutation in the *pssS* gene, which encodes glycosyltransferase belonging to the GTB-type superfamily (GT family 1, CAZy database). Glucose (Glc) is a dominant sugar component of EPS in *R. leguminosarum*, which is present as the first, fourth, fifth, sixth, and seventh residues in

the repeating units (Robertsen et al. 1981). All sugar residues in the unit are bound by β -linkages (β -1,3, β -1,4, and β -1,6); the only exception is Glc at the fourth position, which is linked by an α -1,4 glycosidic bond. Based on amino acid sequence homology, the PssS protein might be involved in the fourth step of the unit synthesis. Similarly, Ivashina and Ksenzenko (2012) have recently proposed a model for the EPS repeating unit assembly in *R. leguminosarum* VF39 and TA1 strains, in which the fourth sugar residue (Glc) is linked by PssS, whereas addition of three further Glc residues is conducted by PssF, PssI/PssG, and PssI/PssH proteins, respectively, which belong to the GTA-type superfamily.

The phenotype of the Rt770 mutant obtained by us confirms that the PssS enzyme plays an essential role in EPS synthesis and symbiosis (Tables 4 and 5). In contrast to the wild-type strain, this EPS-non-producing mutant was unable to establish effective symbiosis with clover; the nodules induced on the host roots were very small, white, and contained only sporadically infected cells, in which bacteroids degenerated prematurely (Figs. 4, 5, and 6). The infection of plant cells was abnormal, visible as “explosion-like” mass endocytosis of the bacteria into the cell cytoplasm. This type of simultaneous endocytosis of a huge number of bacterial cells was previously reported for the *pssD* mutant of *R. leguminosarum* (Król et al. 1998) and the SGEFix⁻1 mutant in pea with a mutation in *sym40* controlling early nodule developmental stages (Tsyganov et al. 1998).

Previously, it was indicated for other EPS-deficient strains of *R. leguminosarum* and *S. meliloti* that absence of EPS led to similar strong disturbances in symbiosis (Rolfe et al. 1996; Guerreiro et al. 2000; Cheng and Walker 1998). Mutations in the *pssA* and *pssD* genes of *R. leguminosarum* bvs. *trifolii* and *viciae* also totally abolished EPS synthesis and resulted in inefficient symbiosis with the compatible host plants, which form indeterminate-type nodules (sporadically formed nodules were small and hardly occupied by bacteria) (Ivashina et al. 1994; Rolfe et al. 1996; van Workum et al. 1997; Król et al. 1998; Laus et al. 2005; Janczarek and Rachwał 2013). The formation of infection threads was only sporadically initiated and they were early aborted. Furthermore, Guerreiro et al. (2000) have reported that mutations in these genes had pleiotropic effects and they, among others, resulted in changed levels of synthesis of several proteins (over 20 proteins in the *pssA* mutant and 8 proteins in the *pssD* mutant). It was established that the symbiotic interaction of the

pssD mutant with its host, clover was impaired in the step of endocytosis of the bacteria into the plant cell cytoplasm (Król et al. 1998). In contrast, an EPS-deficient *pssA* mutant of *R. leguminosarum* bv. *phaseoli* induced nitrogen-fixing nodules on its host, *Phaseolus*, which forms determinate-type nodules (Borthakur et al. 1988; Latchford et al. 1991). This indicates that the significance of EPS in symbiosis is mainly dependent on the kind of the host plant. Similarly, *S. meliloti* strains having mutations in genes encoding glycosyltransferases (*exoY*, *exoA*, *exoL*, *exoM*, *exoO* and *exoW*) were defective in production of succinoglycan and nodule invasion (Reuber et al. 1991; Becker et al. 1993a, b; Cheng and Walker 1998). Also, a lack of the PssV protein responsible for addition of pyruvyl residues to the units abolished production of this polysaccharide and negatively affected symbiosis with alfalfa (Glucksmann et al. 1993; Becker et al. 1993a). Furthermore, mutations in genes involved in the synthesis of nucleotide sugar precursors also caused strong negative effects on both EPS production and infection host plants. For example, a mutation in *R. leguminosarum* *exo5* encoding UDP-glucose dehydrogenase responsible for oxidation of UDP-glucose to UDP-glucuronic acid, affected all glucuronic and galacturonic acid-containing polysaccharides including EPS and, in a consequence, successful colonization of the host roots (Laus et al. 2004). Also, mutations in *S. meliloti* *exoC* and *exoB* genes responsible for the synthesis of glucose-1-phosphate and UDP-galactose, respectively, abolished production of succinoglycan and other polysaccharides containing galactose (galactoglucan and LPS) (Glucksmann et al. 1993; Becker et al. 1993a, b). All these data indicate that EPS is indispensable for host root infection and effective nitrogen-fixation in symbiotic interactions with legumes forming indeterminate nodules. Mutations in genes encoding enzymes involved in the sugar precursor synthesis as well as those encoding glycosyltransferases result in similar effects, i.e., a lack of EPS synthesis and inefficient infection of the host plants.

Several functions are ascribed to rhizobial EPS in symbiosis such as facilitation of bacterial adhesion to and invasion of host roots, a structural function as a component of matrix inside infection threads, and protection against plant defense reactions (Downie 2010; Cheng and Walker 1998; Jaszek et al. 2014). D'Haeze et al. (2004) indicated for *Azorhizobium caulinodans* that a mutant strain producing significantly less EPS that the wild-type bacterium was more sensitive to

oxidative stress and induced only ineffective pseudonodules on its host plant, *Sesbania rostrata*. In addition, these researchers showed a negative correlation between the amount of produced EPS and the level of H₂O₂ inside the bacterial cells. These data indicated that EPS produced by wild-type rhizobia forms a diffusion barrier protecting bacterial cells against huge amounts of hydrogen peroxide encountered during early stages of symbiosis (i.e., legume infection and nodule development). Also, EPS-deficient mutants of *R. leguminosarum* bv. *trifolii* showed decreased survival and adaptation to oxidative stress conditions (Jaszek et al. 2014).

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